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(54) Title: EXTENDED RELEASE ORAL DOSAGE COMPOSITION

(57) Abstract: A compressed bilayer solid composition comprising (a) an immediate release first layer comprising an anti-allergic effective amount of desloratedine and at least one pharmaceutically acceptable excipient and (b) a sustained release second layer comprising an effective amount of a nasal decongestant and a pharmaceutically acceptable sustained release agent wherein the composition contains less than about 2 % of desloratedine decomposition products is disclosed.

EXTENDED RELEASE ORAL DOSAGE COMPOSITION BACKGROUND OF THE INVENTION

This invention relates to a bilayer sustained release oral dosage composition containing a nasal decongestant, e.g., pseudoephedrine in one layer and the non-sedating antihistamine, desloratedine in a second layer and having less than about 2% of desloratedine degradation products. The oral dosage compositions of this invention are useful for treating patients showing the signs and symptoms associated with allergic and/or inflammatory conditions such as the common cold, as well as signs and symptoms associated with allergic and/or inflammatory conditions of the skin and airway passages such as dermatitis, allergic rhinitis, seasonal allergic rhinitis and nasal congestion, upper respiratory diseases, allergic rhinitis and nasal congestion.

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Desloratadine, also called descarbethoxyloratadine, is disclosed in US Patent No. 4,659,716 as a non-sedating antihistamine useful as an anti-allergy agent. US Patent No. 5,595,997 discloses methods and compositions for treating seasonal allergic rhinitis symptoms using desloratadine.

- U. S. Patent Nos. 4,990,535 and 5,100,675 disclose a twice-a-day sustained release coated tablet wherein the tablet coating comprises descarbethoxyloratadine and a hydrophilic polymer and polyethylene glycol, and the tablet core comprises acetaminophen, pseudoephedrine or a salt thereof, a swellable hydrophilic polymer and pharmaceutically acceptable excipients.
- U. S. Patent No. 5,314,697 discloses an extended release tablet containing matrix core comprising pseudoephedrine sulfate and a coating comprising loratedine.

None of the prior art discloses the twice-a-day non-film-coated oral dosage composition of this invention.

The successful development of a formulation of a desloratedinepseudoephedrine twice-a-day product would be desirable, but would require (1) achieving a release rate profile for pseudoephedrine component over an extended period of about twelve hours while maintaining the safety and effectiveness of desloratedine, and (2) minimizing impurity formation due to the interaction between desloratedine and excipients in the pseudoephedrine layer that are incompatible with desloratedine.

It would be desirable for increased patient compliance to have a stable, extended release desloratedine-pseudoephedrine product substantially free of desloratedine impurities and additional polymorphic forms that is effective and safe when used on a twice-a-day or once-a-day basis for the treatment, management and/or mitigation of the signs and symptoms associated with the common cold, as well as allergic and/or inflammatory conditions of the skin or upper and lower airway passages such as seasonal, allergic rhinitis and nasal congestion.

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SUMMARY OF THE INVENTION

We have found that desloratedine discolors and decomposes in the presence of excipients disclosed in the prior art. We have discovered that these problems are substantially solved (a) when the use of an acidic excipient in the desloratedine layer is avoided and when desloratedine is combined with a pharmaceutically acceptable carrier medium comprising a desloratedine protective amount of a pharmaceutically acceptable basic salt, or (b) when a desloratedine-protective amount of a pharmaceutically acceptable antioxidant is present in at least one layer and preferably at least one of said antioxidants is present in each layer of the bilayer tablet.

Thus, this invention provides a compressed bilayer solid composition comprising (1) an immediate release first layer comprising an anti-allergic effective amount of desloratedine and a desloratedine-protective amount of a pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt, or of a desloratedine-protective amount of at least one pharmaceutically acceptable antioxidant; and (2) a sustained release second layer comprising an effective amount of pseudoephedrine or a salt thereof, and a pharmaceutically acceptable sustained release agent, and optionally a desloratedine-protective amount of a pharmaceutically acceptable antioxidant.

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Thus, in one preferred embodiment, this invention provides a compressed bilayer solid composition comprising (1) one layer- an immediate release first layer-comprising an anti-allergic effective amount of desloratedine and desloratedine-protective amount of a pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt, and (2) another layer-a sustained release second layer- comprising an effective amount of pseudoephedrine or a salt thereof, and a pharmaceutically acceptable sustained release agent.

The pharmaceutical compositions of the present invention contain less than about 2.0% of desloratadine decomposition products such as N-formyldesloratadine (see Chart I) when such compositions are stored at 25°C and about 60% relative humidity for extended time periods, e.g., about 18 months.

In a preferred embodiment, this invention provides a compressed bilayer solid composition comprising:

(a) an immediate release first layer comprising:

	INGREDIENT	mg/composition
	Desloratadine, micronized	2.5
	Corn starch	11.0
	Dibasic calcium phosphate dihydrate	53.0
20	Microcrystalline cellulose	30.22
	Talc	3.0
	FD&C Blue dye No. 2 Aluminium Lake 5627	0.28
	TOTAL IN FIRST LAYER	100.00

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(b) a second sustained release second layer comprising:

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	INGREDIENT	mg/composition
	Pseudoephedrine Sulfate	120.0
5	Hydroxypropyl Methylcellulose	105.0
	Microcrystalline cellulose	100.0
	Povidone	18.0
	Silicon Dioxide	5.0
	Magnesium stearate	2.0
10	TOTAL IN SECOND LAYE	R 350.0

The above-listed preferred compressed bilayer composition contains less than about 2.0% of desloratedine decomposition products such as N-formyl-desloratedine (see Chart I) when such compositions are stored at 25°C and about 60% relative humidity for extended time periods of about 18 months.

Thus, in another preferred embodiment, the present invention also provides a compressed bilayer solid composition comprising (1) an immediate release first layer comprising an anti-allergic effective amount of desloratedine and a desloratedine-protective amount of at least one pharmaceutically acceptable antioxidant; and (2) a sustained release second layer comprising an effective amount of pseudoephedrine or a salt thereof, a pharmaceutically acceptable sustained release agent, and a desloratedine-protective amount of a pharmaceutically acceptable antioxidant. The above-listed preferred compressed bilayer composition contains less than about 2.0% of desloratedine decomposition products such as N-formyldesloratedine (see Chart I) when such compositions are stored at 25°C and about 60% relative humidity for extended time periods of about 18 months.

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The present invention provides a compressed bilayer solid composition comprising (a) an immediate release first layer comprising an anti-allergic effective

amount of desloratadine and at least one pharmaceutically acceptable excipient and (b) a sustained release second layer comprising an effective amount of a nasal decongestant and a pharmaceutically acceptable sustained release agent. In a preferred embodiment, the compressed bilayer solid composition contains less than about 2.0% of desloratadine decomposition products such as N-formyl desloratadine after storage for about 18 months, and wherein at least about 80% of the desloratadine dissolves in 0.1N HCl at 37°C in about 45 minutes.

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In another preferred embodiment, the present invention also provides a compressed bilayer solid composition comprising (1) an immediate release first layer comprising 5 mg of desloratedine and desloratedine-protective amount of a pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt ,and (2) a sustained release second layer comprising 120 mg of pseudoephedrine sulfate, and a pharmaceutically acceptable sustained release agent. This preferred composition provides a 24-hr dose of desloratedine and a 12-hr dose of pseudoephedrine sulfate.

Thus, the present invention also provides a method of treating and/or preventing allergic and inflammatory conditions of the upper and lower airway passages and skin which comprises administering to a patient in need of such treating an effective amount of a compressed bilayer solid composition of the present invention.

DETAILED DESCRIPTION OF THE INVENTION

During the course of development of the compositions of the present invention, deslorated was found to be unstable and to discolor when stored in combination with various excipients such as those disclosed in U.S. Patent No. 5,314,697 as part of the matrix core containing pseudoephedrine sulfate. The excipients causing discoloration and instability of deslorated include acidic excipients having a pH of less than 7 in water such as organic acids, such as stearic acid, povidone, crospovidone as well as the hydroxycarboxylic acid, ascorbic acid, and carbonyl-containing materials such as lactose, and ethylcellulose and hydroxylpropyl methylcellulose. Binders like povidone and polymers such as hydroxypropyl methylcellulose are useful as a polymer matrix for

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the sustained release of the pseudoephedrine sulfate from the inner polymer matrix core.

We also discovered that metal ions catalyzed were involved in the formation of desloratadine degradation products.

We have discovered two solutions to inhibit and/or prevent formation of desloratedine degradation products. In one preferred embodiment, a desloratedine-protective amount of a pharmaceutically acceptable anti-oxidant should be present in at least one of the bilayers, preferably one of said antioxidant in each layer.

In a second preferred embodiment, we also discovered that it is possible to prepare a bilayer tablet containing desloratedine in an immediate release first layer in intimate contact with a sustained release second layer containing a nasal decongestant and excipients incompatible with desloratedine by incorporating a desloratedine protective amount of a pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt into the immediate release desloratedine layer.

The term " in intimate contact" as used herein in reference to the two layer forming the bilayer tablet means that there is with no film interface between the two layers.

The term "pharmaceutically acceptable antioxidant" as used herein in reference to desloratedine (formula I in the Chart) means a pharmaceutically acceptable chelating agent that protects desloratedine from the formation of degradation products including, but not limited to those of the formulas II-V listed in the Chart ,e.g.,N-formyl-desloratedine or N-formyl DL(formula II in the Chart), N-hydroxylamine of DL (formula V in the Chart) N-oxide of DL(formula IV in the Chart), and the 3'-hydroxyl N-oxide of DL(formula III in the Chart). The structures listed in the Chart were determined by standard physiochemical techniques, e.g., LC-MS, and LC-NMR.

Typically suitable pharmaceutically acceptable antioxidants for DL are pharmaceutically acceptable chelating agents such as those disclosed in "Chelating Agents", pages 764-794, Vol. 5 of KIRTH-OTHMER, ENCYCLOPEDIA OF CHEMICAL TECHNOLOGY, 4th Edition, 1993, John Wiley & Sons Inc., NY, and preferably including, but not limited to, hydroxycarboxylic acids, such as tartaric acid, citric acid and gluconic acid, and pharmaceutically acceptable salts thereof, aminocarboxylic acids such as edetic acid (ethylenediamine tetraacetic acid) and pharmaceutically acceptable

salts thereof such as edetate calcium disodium, edetate disodium, edetate trisodium, and edetate tetrasodium. Edetate disodium and citric acid are the preferred pharmaceutically acceptable antioxidants. Use of the hydroxycarboxylic acid, ascorbic acid, is to be avoided

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The desionatedine protective amount of a pharmaceutically acceptable antioxidant in the DL immediate release layer is in the range of about 0.1% to about 10% by weight, preferably about 1% to 8% or about 1% to about 6%, more preferably about 4% to about 8%, or about 4% to about 6%, or most preferably about 5% to about 6%. The desloratadine protective amount of a pharmaceutically acceptable antioxidant in the PES sustained release layer is in the range of 0% to about 10%, preferably about 0.1% to about 10%, or about 0.1% to about 3%, more preferably about 1 to about 2%, and most more preferably about 1.0%. In a preferred embodiment of the present invention, about 1.0% by weight of a pharmaceutically acceptable antioxidant, e.g., edetate disodium, is present in the PES sustained release layer. In another preferred embodiment, about 6% by weight of a mixture of two pharmaceutically acceptable antioxidants, e.g., edetate disodium and citric acid, are present in the DL immediate release layer in a ratio of about 5:1 to about 1:5, preferably about 5:1, and about 1% of a pharmaceutically acceptable antioxidant, e.g., edetate disodium, is present in the sustained release layer. In another preferred embodiment, about 5% by weight of one pharmaceutically acceptable antioxidant, e.g., edetate disodium, is present in the DL immediate release layer.

In other preferred embodiments, about 5.0 mg (a 24-hour supply) of DL is present in the DL immediate release layer, and 120 mg (a 12-hour supply) of the nasal decongestant pseudoephedrine sulfate is present in the sustained release layer(see Examples 4,5&6). In one preferred embodiment, the dibasic phosphate salt preferably dibasic calcium phosphate dihydrate is present in the DL immediate release layer and no pharmaceutically acceptable antioxidant is present in either layer (see Example 4). In another preferred embodiment, 5.0 mg (a 24-hour supply) of DL and about 0.1 to about 10% of at least one antioxidant is present in the DL immediate release layer, preferably about 4% to about 6% of a mixture of two antioxidants, e.g., edetate disodium and citric acid, in a ratio of 5:1 to 1:1, preferably in a ratio of 5:1, and about 0.1% to about 10% preferably about 0.1% to about 5%, more preferably about 0.1% to

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about 3%, most more preferably about 1.0% of an antioxidant, e.g., edetate disodium, is present in the PES sustained release layer(see Examples 5&6).

The desloratedine was found to have an acceptable immediate release profile from the second layer (80% release in 0.1N HCl in less than about 45 min.) and contain less than about 2% of desloratedine degration products even after storage for at least 18 months at 25° C and about 60% relative humidity ("RH").

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The phrase "allergic and inflammatory conditions of the skin and airway passages" means those allergic and inflammatory conditions and symptoms found on the skin and in the upper and lower airway passages from the nose to the lungs. Typical allergic and inflammatory conditions of the skin and upper and lower airway passages include seasonal and perennial allergic rhinitis, non-allergic rhinitis, asthma including allergic and non-allergic asthma, sinusitis, colds (in combination with a NSAID, e.g., aspirin ibuprofen or APAP) and/or a decongestant e.g. pseudoephedrine), dermatitis, especially allergic and atopic dermatitis, and urticaria and symptomatic dermographism as well as retinophathy, and small verssel diseases, associated with diabetes mellitus.

The amount of desloratadine effective for treating or preventing allergic and inflammatory conditions of the skin and upper and lower airway passages will vary with the age, sex, body weight and severity of the allergic and inflammatory condition of the patient. Typically, the amount of desloratadine effective for treating or preventing such allergic and inflammatory conditions is in the range of about 2.5 mg/day to about 60 mg/day, preferably about 2.5 mg/day to about 20 mg/day, or about 4.0 mg/day to about 15 mg/day, or about 5.0 mg/day to about 10 mg/day, more preferably about 5.0 mg/day to about 10.0 mg/day, and most preferably about 5.0 mg/day in one dose or in two divided doses of 2.5 mg/dose.

Desloratadine is a non-sedating long acting histamine antagonist with potent selective peripheral H1-receptor antagonist activity. Following oral administration, loratadine is rapidly metabolized to descarboethoxyloratadie or desloratadine, a pharmacologically active metabolite. *In vitro* and *in vivo* animal pharmacology studies have been conducted to assess various pharmacodynamic effects of desloratadine and loratadine. In assessing antihistamine activity in mice

(comparison of ED₅₀ value), desloratedine was relatively free of producing alterations in behavior alterations in behavior, neurologic or autonomic function. The potential for desloratedine or loratedine to occupy brain H1-receptors was assessed in guinea pigs following i.p. administration and results suggest poor access to central histamine receptors for desloratedine or loratedine.

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In addition to antihistaminic activity, deslorated has demonstrated antiallergic and anti-inflammatory activity from numerous *in vitro* and *in vivo* tests.

These *in vitro* tests (mainly conducted on cells of human origin) have shown that desloratedine can inhibit many events in the cascade of allergic inflammation.

These anti-inflammatory effects for desloratedine are independent of the H1-

- antagonist effect of desloratadine and include:
 The release of inflammatory mediators histamine, truptase, leukotriene and prostaglandin D2 from mast cells;
- The release of inflammatory cytokines including IL-4, IL-6, IL-8 and IL-13;
- The release of the inflammatory chemokines such as RANTES (regulated upon activation, normal T cell expressed and presumably secreted);
 - Superoxide anion production of polymorphonuclear neutrophils;
 - The expression of cell adhesion molecules such as intracellular adhesion molecules (ICAM-1) and P-selection in endothelial cells; and
- Eosinophil migration and adhesion
 In vivo studies also suggest that an inhibitory effect of desloratadine on allergic bronchospasm and cough can also be expected.

The clinical efficacy and safety of deslorated has been documented in over 3,200 seasonal allergic rhinitis patients in 4 double-blind, randomized clinical trials. The results of these chemical studies demonstrated the efficacy of deslorated in the treatment of adult and adolescent patients with seasonal rhinitis.

The nasal decongestants useful in the present invention include phenylpropanolamine, phenylephrine and and pseudoephedrine. Pseudoephedrine as well as pharmaceutically acceptable acid additional salts, e.g., those of HCl or H₂SO₄, is a sympathomimetic drug recognized by those skilled in the art as a safe therapeutic agent effective for treating nasal congestion and is commonly administered orally and

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concomitantly with an antihistamine for treatment of nasal congestion associated with allergic rhinitis. The use of pseudoephedrine as a nasal decongestant in the present invention is preferred; the use of about 120 mg pseudoephedrine sulfate in the extended release layer is more preferred.

In the course of development of the compressed bilayer oral dosage composition of this invention, it was discovered that the selection of the polymers for the extended release layer was critical to achieve the desired extended release period of at least 12 hours, for pseudoephedrine sulfate. For example, the use of hydroxypropyl methylcellulose 4,000 cps or 15,000 cps as polymers in the matrix core did not provide this more preferred extended release period of at least 16 hours for dose of pseudoephedrine sulfate. We discovered that only by selecting for inclusion into the matrix core of specific weight ratios of three specific polymers was the desired pseudoephedrine release profile achieved. Only by combining (1) about one part by weight, preferably 1.05 parts by weight of hydroxypropyl methylcellulose 2208 USP, 100,000 cps with (2) about one part by weight, preferably 1.0 parts by weight of microcrystalline cycllulose together with (3) about 0.15-0.20 part by weight., preferably 0.17-0.18 parts by weight of povidone (per 1.05 parts by weight of hydroxypropyl methylcellulose) as a secondary binder was the more preferred extended release profile of at least 12 hours for pseudoephedrine sulfate from the extended release layer. The extended release layer also contains specific amounts of silicon dioxide as a glidant and magnesium stearate as a lubricant. The tablet hardness 20 ± 4 Strong-Cobb Units (SCU) is not greatly affected by the higher level of lubricant (6mg/tablet) but it is preferred to maintain the lubricant level at 1/9 part by weight of lubricant to one part by weight of povidone as secondary binder.

The term "lubricant' as used herein refers to a substance added to the dosage form to enable the dosage form, e.g., a tablet, after it has been compressed to release from the mold or die.

Suitable lubricants include talc, magnesium stearate, calcium stearate, stearic acid, hydrogenated vegetable oils and the like. Preferably, magnesium stearate or talc is used.

The term "glidants" as used herein refers to a substance, such as an anti-caking agent, which improves the flow characteristics of a powder mixture.

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Suitable glidants include silicon dioxide and talc. Preferably, silicon dioxide is used.

The term "binders" as used herein means any material that is added to pharmaceutical compositions to help hold such compositions together and release the medicament therefrom.

Suitable binders are selected those found in NF XVIII, page 2206 (1995) and include povidones, starches, celluloses, alginates, and gums and low molecular weight hydroxypropyl melthyl celluloses, especially hydroxypropyl methyl cellulose 2910.

The term "pharmaceutically acceptable water insoluble basic calcium, magnesium and aluminium salts" as used herein means the pharmaceutically acceptable carbonates, phosphates, silicates and sulfates of calcium, magnesium and aluminum or mixtures thereof. Typically suitable pharmaceutically acceptable basic salts include calcium sulfate anhydrous, hydrates of calcium sulfate, such as calcium sulfate dihydrate, magnesium sulfate anhydrous, hydrates of magnesium sulfate, dibasic calcium phosphate, dibasic calcium silicate, magnesium trisilicate, magnesium phosphate, aluminum silicate, and hydrates of magnesium phosphate, aluminum phosphate is more preferred. The use of dibasic calcium phosphate dihydrate is most preferred.

The desloratedine-protective amount of a pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt is in the range of about 50-60% of the DL immediate release layer, and the w/w ratio of the pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt to DL is in the range of about 8:1 to about 40:1, more preferably is in the range of about 10:1 to about 20:1, and most preferably is in the range of about 10:1 to about 11:1.

In the preferred embodiment of the present invention wherein a desloratadine protective amount of a pharmaceutically acceptable antioxidant is present, the water insoluble basic calcium salt is not present in the immediate release layer containing desloratadine; in its place, at least one, preferably two pharmaceutically acceptable antioxidants are present, e.g., edetate sodium and citric acid and the amount of microcrystalline cellulose is increased. In addition, when the pharmaceutically acceptable antioxidant is used in place of the water insoluble basic calcium, magnesium or aluminum salt, the povidone in the sustained release layer is replaced by another

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binder, preferably a low molecular weight hydroxypropyl methyl cellulose ("HPMC"), preferably HPMC 2910.

The oral dosage composition of this invention also provides a shelf life of up to 18 months so long as the tablets are stored in standard package at between 2° and 30° C in an ambient environment of 60% relative humidity.

In the preparation of the bilayer tablet, the sustained release layer is compacted first. The immediate release second layer is added on top and a compression force sufficient to form a bilayer tablet is applied in the range of 8-12 kilo Newtons, preferably about 9 kilo Newtons(kN).

The dried sustained release granulation is milled and blended with requisite amounts of silicon dioxide and magnesium stearate. In a preferred embodiment, a pharmaceutically acceptable blue dye containing EDTA as a chelating agent is incorporated into the immediate release deslorated in layer. Use of a pharmaceutically acceptable blue dye, e.g. FD& C blue dye No. 2 Aluminum Lake 5627 is preferred.

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EXAMPLE I

This example illustrates preparation of the preferred oral dosage composition of this invention. The ingredients and specific amounts thereof are listed below.

A. Method of Manufacture of the Immediate Release Layer

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 Prepare starch paste by dispersing the paste portion of corn starch into purified water in a suitable container equipped with an agitator.

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2. While mixing, heat the contents to approximately 95°C and maintain the temperature for approximately 30 minutes.

3. After Step 2 is completed, add an additional purified water and allow the starch paste to cool to approximately 50°C.

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4. While mixing, add desloratedine to the starch paste. Continue mixing during the granulation step.

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- Pass the FD&C blue No. 2 aluminum lake containing EDTA as a chelating,e.g., Spectra Spray Med Blue, with the required amount of dibasic calcium phosphate through a suitable sieve or mill.
- 6. Charge to a suitable fluid bed processing bowl the remaining dibasic calcium phosphate dihydrate, the milled material from Step 5, a portion of the corn starch, and a portion of microcrystalline cellulose. Place the processing bowl into the fluid bed processor.
- 7. Fluidize the powder bed until the product temperature reaches approximately 29°C.
 - 8. Begin granulating the powder by pumping the starch paste from Step 4 into the fluidized bed at a suitable spray rate and a bed temperature of approximately 22°C.
 - 9. Continue to dry the granulation at an inlet air temperature of approximately 60°C until a final loss on drying (LOD) of 2% or less is achieved.
 - 10. Pass the dried granulation through a suitable sieve or mill.
 - 11. Charge the granulation to a suitable blender and add the requisite amounts of the remaining portion of microcrystalline cellulose, corn starch, and talc. Blend for 5 minutes.

B. Manufacture of Sustained Release Mix:

 Charge purified water and alcohol to a suitable container equipped with an agitator.

- 2. Dissolve povidone in the water/alcohol mixture. Continue mixing for a minimum of 10 minutes.
- Mix hydroxypropyl methylcellulose, pseudoephedrine sulfate and
 microcrystalline cellulose in a suitable granulator.
 - Granulate the mix with the povidone solution, using additional water/alcohol mixture if necessary to achieve the appropriate granulation consistency.
 - 5. Dry the wet granulation at approximately 50°C in a suitable dryer until the loss on drying (LOD) is between 1% and 3%.
 - 6. Pass the dried granulation through a suitable sieve or mill.
 - 7. Charge the milled granulation to a suitable blender.
 - 8. Pass the silicon dioxide through a No. 30 mesh screen into the blender.
- Blend the requisite amount of screened silicon dioxide with the granulation for approximately 10 minutes in a suitable blender.
 - 10. Pass the magnesium stearate through a No. 30 mesh screen.
- 25 11. Blend the requisite amount of screened magnesium stearate with the mix from Step 9 for 5 minutes.

C. Compression:

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Compress the two blends to specifications as bilayer tablets using a suitable double-layer tablet press using a compression force of 9k Newtons. Compress the sustained release layer first.

Tablet Weight: 450 mg± 5%

- Sustained release layer: 350 mg± 5%

- Immediate release layer: 100 mg ± 5%

Hardness: 20 ± 4 SCU (Strong Cobb units)

The following bilayer tablet was prepared using the above procedure.

10 <u>Desloratadine Immediate Release Layer:</u>

	INGREDIENT	mg/composition
	Desloratadine, micronized	2.5
15	Corn Starch NF/Ph.Eur.	11.0
	Dibasic Calcium Phosphate Dihydrate USP/Ph.	Eur. 53.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	30.22
	Talc USP/Ph.Eur.	3.0
	Dye FD&C Blue No. 2 Aluminium Lake 5627	0.28
20	Water Purified USP/Ph.Eur.	
	TOTA	L 100.00
	and	

Pseudoephedrine Sulfate Sustained Release Layer

25	INGREDIENT	mg/composition
	Pseudoephedrine Sulfate USP	120.0
	Hydroxypropyl Methylcellulose USP/Ph.Eur.	105.0
	Microcrystalline Cellulose 2208,	
30	100,000cpsNF/Ph.Eur./JP	100.0
	Povidone USP/Ph.Eur./JP	18.0
	Silicon Dioxide NF	5.0

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	Magnesium Stearate NF/Ph.Eur.JP(Non-Bovine)	2.0
	Water Purified USP/Ph.Eur.	
	Alcohol USP/3A Alcohol	
	TOTAL	350.0
5	TOTAL TABLET	450.0

Hardness:

20 ± 4 SCU (Strong Cobb units)

EXAMPLE 2

The procedure of Example 1 was used; edetate disodium was used in place
of the dibasic calcium salt and the amount of microcrystalline cellulose in the DL
layer was increased. Edetate disodium was used in the sustained release layer and
hydroxypropyl methylcellulose 2910 was used in place of povidone.

Desloratadine Immediate Release Layer:

15	INGREDIENT	mg/composition
	Desloratadine, micronized	2.5
	Corn Starch NF/Ph.Eur.	8.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	71.35
20	Edetate Disodium	5.0
	Talc USP/Ph.Eur.	3.0
	Dye FD&C Blue No. 2 Aluminium Lake 5627	0.15
	Water Purified USP/Ph.Eur.	
	TOTA	L 100.00
25	and	

Pseudoephedrine Sulfate Sustained Release Layer

	INGREDIENT	mg/composition
30	Pseudoephedrine Sulfate USP	120.0
•	Hydroxypropyl Methylcellulose 2208,USP/Ph.Eu	
	Microcrystalline Cellulose NF/Ph.Eur./JP	103.5

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	Edetate Disodium	3.5
	Hydroxypropyl Methylcellulose 2910 USP/Ph.Eur./JP	10.5
	Silicon Dioxide NF	5.0
	Magnesium Stearate NF/Ph.Eur.JP(Non-Bovine)	2.5
5	Water Purified USP/Ph.Eur.	
	Alcohol USP/3A Alcohol	
	TOTAL	350.0
	TOTAL TABLET	450.0

10 Hardness:

20 ± 4 SCU (Strong Cobb units)

EXAMPLE 3

The procedure of Example 2 was used, but I mg of citric acid was added to the DL layer and .the amount of microcrystalline cellulose was decreased by 1 mg.

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Desloratadine Immediate Release Layer:

	INGREDIENT	mg/composition
	Desloratadine, micronized	2.5
20	Corn Starch NF/Ph.Eur.	18.0
	Edetate Disodium	5.0
	Citric Acid	1.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	70.35
	Talc USP/Ph.Eur.	3.0
25	Dye FD&C Blue No. 2 Aluminium Lake 5627	0.15
	Water Purified USP/Ph.Eur.	
	TOTA	L 100.00

And Pseudoephedrine Sulfate Sustained Release Layer

30	INGREDIENT	mg/composition
	Pseudoephedrine Sulfate USP	120.0
	Hydroxypropyl Methylcellulose 2208,100,000cp	os .

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	USP/Ph.Eur.	105.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	103.5
	Edetate Disodium	3.5
	Hydroxypropyl Methylcellulose 2910	10.5
5	Silicon Dioxide NF	5.0
	Magnesium Stearate NF/Ph.Eur.JP(Non-Bovine)	2.5
	Water Purified USP/Ph.Eur.	
	Alcohol USP/3A Alcohol	
	TOTAL	350.0
10	TOTAL TABLET	450.0

Hardness:

20 ± 4 SCU (Strong Cobb units)

EXAMPLE 4

The procedure of Example 1 was used. The bilayer tablet of Example 1 was modified by including 5.0 mg of desloratedine in the immediate release layer-(a 24 hour dose)-with the appropriate changes in amounts of the other ingredients and using the 12-hr dose pseudoephedrine sustained release layer of Example 1.

Hardness:20 ± 4 SCU (Strong Cobb units)

Desloratadine Immediate Release Layer:

20	INGREDIENT	mg/composition
	Desloratadine, micronized	5.0
	Corn Starch NF/Ph.Eur.	11.0
	Dibasic Calcium Phosphate Dihydrate USP/Ph.	Eur. 53.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	27.72
25	Talc USP/Ph.Eur.	3.0
	Dye FD&C Blue No. 2 Aluminium Lake 5627	0.28
	Water Purified USP/Ph.Eur.	
	TOTA	L 100.00
	and	

30 <u>Pseudoephedrine Sulfate Sustained Release Layer</u>

INGREDIENT mg/composition

120.0

Pseudoephedrine Sulfate USP

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Hydroxypropyl Methylcellulose 2208,1000,00cps	
USP/Ph.Eur.	105.0
Microcrystalline Cellulose NF/Ph.Eur./JP	100.0
Povidone USP/Ph.Eur./JP	18.0
5 Silicon Dioxide NF	5.0
Magnesium Stearate NF/Ph.Eur.JP(Non-Bovine)	2.0
Water Purified USP/Ph.Eur.	
Alcohol USP/3A Alcohol	
TOTAL	350.0
10 TOTAL TABLET	450.0

EXAMPLE 5

The procedure of Example 1 was used and the bilayer tablet of Example 4 was modified by replacing the dibasic calcium phosphate dihydrate in the immediate release layer with 10 mg of edetate disodium and increasing the amount of microcrystalline cellulose by 2.7 mg. Hardness:20 ± 4 SCU (Strong Cobb units)

Desloratadine Immediate Release Layer:

	INGREDIENT	mg/composition
	Desloratadine, micronized	5.0
20	Corn Starch NF/Ph.Eur.	36.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	142.7
	Edetate Disodium	10.0
•	Talc USP/Ph.Eur.	6.0
	Dye FD&C Blue No. 2 Aluminium Lake 5627	0.30
25	Water Purified USP/Ph.Eur.	
	TOTA	L 200.00

and

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	Pseudoephedrine Sulfate Sustained Release Layer	
	INGREDIENT	mg/composition
30	Pseudoephedrine Sulfate USP	120.0
	Hydroxypropyl Methylcellulose 2208,1000,00cp	os
	USP/Ph.Eur.	105.0

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	Microcrystalline Cellulose NF/Ph.Eur./JP	103.5
	Hydroxypropyl Methylcellulose 2910	10.5
	Edetate Disodium	3.5
	Silicon Dioxide NF	5.0
5	Magnesium Stearate NF/Ph.Eur.JP(Non-Bovine)	2.5
	Water Purified USP/Ph.Eur.	
	Alcohol USP/3A Alcohol	
	TOTAL	350.0
10	TOTAL Tablet Weight	550.0

EXAMPLE 6

The bilayer tablet of Example 5 was modified by adding 2.0 mg of citric acid to the immediate release layer and decreasing the amount of microcrystalline cellulose by 2.7 mg and using the pseudoephedrine sustained release layer of Example 1. Hardness: 20 ± 4 SCU (Strong Cobb units)

Desloratadine Immediate Release Layer:

	INGREDIENT	mg/composition
	Desloratadine, micronized	5.0
20	Corn Starch NF/Ph.Eur.	36.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	140.7
	Edetate Disodium	10.0
	Citric Acid	2.0
	Talc USP/Ph.Eur.	6.0
25	Dye FD&C Blue No. 2 Aluminium Lake 5627	0.30
	Water Purified USP/Ph.Eur.	
	TOTA	L 200.00

And Pseudoephedrine Sulfate Sustained Release Layer

	<u>INGREDIENT</u>	mg/composition
30	Pseudoephedrine Sulfate USP	120.0
	Hydroxypropyl Methylcellulose 2208,1000,00cp	os
	USP/Ph.Eur.	105.0

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	Microcrystalline Cellulose NF/Ph.Eur./JP	103.5
	Hydroxypropyl Methylcellulose 2910	10.5
	Edetate Disodium	3.5
	Silicon Dioxide NF	5.0
5	Magnesium Stearate NF/Ph.Eur.JP(Non-Bovine)	2.5
	Water Purified USP/Ph.Eur.	
	Alcohol USP/3A Alcohol	
	TOTAL	350.0
10	TOTAL Tablet Weight	550.0

The *in vitro* dissolution profile of the tablets of Examples 1-6 were measured in a stirred 0.1N HCl solution at 37°C (1st hour) and thereafter in a stirred phosphate buffer having a pH of 7.5 at 37°C. The 80% of desloratedine in the immediate release layers was dissolved within the first 30 minutes and the total dose of pseudoephedrine sulfate in the sustained release layers was slowly released via erosion and dissolution mechanisms over a period of at least 12 hours.(with 30-45% in 1 hr, 50-605% in 2 hrs. and ≥80% in 6 hrs).

Similar results would be expected if a decongestant effective amount of another pharmaceutically acceptable pseudoephedrine salt, e.g., pseudoephedrine hydrochloride was used in place of pseudoephedrine sulfate.

The compositions of the present invention are useful for treatment of allergic and/or inflammatory conditions of the skin (e.g. urticaria) and the upper and lower airway passages including the nasal and non-nasal symptoms of seasonal allergic rhinitis including nasal congestion in a patient in need of such treating. The precise dosage and dosage regimen may be varied by the attending clinician in view of the teachings herein depending upon the requirements of the patient, e.g., the patient's age, sex and the severity of the allergic and/or inflammatory condition being treated. Determination of the proper dosage and dosage regimen for a particular patient will be within the skill of the attending clinician.

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While we have hereinabove presented a number of preferred embodiments of this invention by way of example, it is apparent that the scope of the invention is to be defined by the scope of the appended claims.

WHAT IS CLAIMED IS:

- A compressed bilayer solid composition comprising (1) a first layer
 comprising an anti-allergic effective amount of desloratedine and a desloratedine-protective amount of a pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt, or of a desloratedine-protective amount of at least one pharmaceutically acceptable antioxidant; and (2) a second layer comprising an effective amount of pseudoephedrine or a salt thereof, and a pharmaceutically acceptable excipient, and optionally a desloratedine-protective amount of a pharmaceutically acceptable antioxidant.
 - 2) A compressed bilayer solid composition comprising (1) a first layer comprising an anti-allergic effective amount of desloratedine and a desloratedine-protective amount of at least one pharmaceutically acceptable antioxidant; and (2) a second layer comprising an effective amount of pseudoephedrine or a salt thereof, a pharmaceutically acceptable excipient, and a desloratedine-protective amount of a pharmaceutically acceptable antioxidant.
- 20 3) A compressed bilayer solid composition comprising (1) a first layer comprising an anti-allergic effective amount of desloratedine and desloratedine-protective amount of a pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt, and (2) a second layer comprising an effective amount of pseudoephedrine or a salt thereof.

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4) A compressed bilayer solid composition comprising (a) an immediate release first layer comprising an anti-allergic effective amount of desloratedine and at least one pharmaceutically acceptable excipient and (b) a sustained release second layer comprising an effective amount of a nasal decongestant and a pharmaceutically acceptable excipient, wherein the total amount of desloratedine degradation products is less than about 2%.

- 5) The compressed bilayer solid composition of any preceding claim wherein the first layer is an immediate layer and wherein the second layer is a sustained release layer containing a pharmaceutically acceptable sustained release agent.
- 5 6) The compressed bilayer solid composition of claim 5 wherein the nasal decongestant is pseudoephedrine, or a pharmaceutically acceptable salt thereof.

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- 7) The compressed bilayer solid composition of any preceding claim wherein at least about 80% of the desloratedine dissolves in a 0.1N HCl solution at 37°C in about 45 minutes.
- 8) The compressed bilayer solid composition of any preceding claim wherein the amount of N-formyldesloratedine is less than about 0.5% after storage at 25°C and 60% relative humidity for an extended time period.
- 9) The compressed bilayer solid composition of claim 1 or 2 wherein about 0.1 % to about 10% of a pharmaceutically acceptable antioxidant is present in each layer.
- 20 10) The compressed bilayer solid composition of any preceding claim wherein the anti-allergic effective amount of desloratedine in the first layer is about 2.5 mg.
 - 11) The compressed bilayer solid composition of any preceding claim wherein the anti-allergic effective amount of desloratedine in the first layer is about 5.0 mg.
 - 12) The compressed bilayer solid composition of claim 1 or 2 wherein two pharmaceutically acceptable antioxidants are present in the desloratedine layer.
- 13) The compressed bilayer solid composition of claim 1 or 3 wherein30 an immediate release first layer comprises:

	INGREDIENT	<u>mg/c</u>	composition
	Desloratadine, micronized		2.5
	Corn Starch		11.0
5	Dibasic Calcium Phosphate Dihydrate		53.0
	Microcrystalline Cellulose		30.22
	Talc		3.0
	Dye FD+C Blue No. 2 Aluminium Lake		0.28
		TOTAL	100.00

10 and

and wherein an sustained release layer comprises

	INGREDIENT	<u>!</u>	ng/compositio	<u>n</u>
	Pseudoephedrine Sulfate		120.0	
15	Hydroxypropyi Methylcellulose		105.0	
	Microcrystalline cellulose		100.0	
	Povidone		18.0	
	Silicon Dioxide		5.0	
	Magnesium stearate		2.0	
20		TOTAL	350.0	

14) The compressed bilayer solid composition of claim 1 or 3 wherein an immediate release first layer comprises:

	INGREDIENT	Ī	mg/composition
25	Desloratadine, micronized		2.5
	Corn Starch		18.0
	Microcrystalline Cellulose		70.35-71.35
	Edetate Disodium		5.0
	Citric Acid		0-1.0
30	Talc		3.0
	Dye FD+C Blue No. 2 Aluminium Lake		0.28
		TOTAL	100.00

and

and wherein an sustained release layer comprises:

5	INGREDIENT	<u>m</u>	ng/composition
	2.4.		420.0
	Pseudoephedrine Sulfate		120.0
	Hydroxypropyl Methylcellulose 2208		105.0
	Microcrystalline cellulose		103.5
10	Edetate Disodium		3.5
	Hydroxypropyl Methylcellulose 2910		10.5
	Silicon Dioxide		5.0
	Magnesium stearate		_2.0
		TOTAL	350.0

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- 15) A compressed bilayer solid composition comprising (1) a first layer comprising 2.5 or 5 mg of desloratedine and desloratedine-protective amount of a pharmaceutically acceptable water insoluble basic calcium, magnesium or aluminum salt, and (2) a second layer comprising 120 mg of pseudoephedrine or a salt thereof, and a pharmaceutically acceptable excipient.
- 16) A compressed bilayer solid composition comprising (1) a first layer comprising 2.5 mg or 5.0 mg of desloratedine and a desloratedine-protective amount of at least one pharmaceutically acceptable antioxidant; and (2) a second layer comprising 120 mg of pseudoephedrine or a salt thereof, a pharmaceutically acceptable excipient, and a desloratedine-protective amount of a pharmaceutically acceptable antioxidant.
- 17) The compressed bilayer solid composition of claim 15 or 16 wherein the amount of desloratedine in the first layer is about 2.5 mg.

- 18) The compressed bilayer solid composition of any preceding claim 15 or 16 wherein the amount of desloratadine in the first layer is about 5.0 mg.
- The compressed bilayer solid composition of claim 1 or 3 wherein
 the immediate release first layer comprises:

Desloratadine Immediate Release Layer:

	INGREDIENT	mg/composition
10	Desloratadine, micronized Corn Starch NF/Ph.Eur.	5.0 11.0
	Dibasic Calcium Phosphate Dihydrate USP/Ph.I	Eur. 53.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	27.72
	Talc USP/Ph.Eur.	3.0
15	Dye FD&C Blue No. 2 Aluminium Lake 5627	0.28
	Water Purified USP/Ph.Eur.	-
	TOTAL	100.00

and

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Pseudoephedrine Sulfate Sustained Release Layer

	INGREDIENT	mg/composition
25	Pseudoephedrine Sulfate USP Hydroxypropyl Methylcellulose 2208,1000,00cp	120.0 s
	USP/Ph.Eur.	105.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	100.0
	Povidone USP/Ph.Eur./JP	18.0
30	Silicon Dioxide NF	5.0
	Magnesium Stearate NF/Ph.Eur.JP(Non-Bovine	e) 2.0
	Water Purified USP/Ph.Eur.	

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Alcohol USP/3A Alcohol		
	TOTAL	350.0
	TOTAL TABLET	450.0

The compressed bilayer solid composition of claim 1 or 2 wherein 20) 5 an immediate release first layer comprises:

Desloratadine Immediate Release Layer:

	INGREDIENT	mg/composition
10		
	Desloratadine, micronized	5.0
	Corn Starch NF/Ph.Eur.	36.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	140.7-142.7
	Edetate Disodium	10.0
15	Citric Acid	0-2.0
	Talc USP/Ph.Eur.	6.0
	Dye FD&C Blue No. 2 Aluminium Lake 5627	0.30
	Water Purified USP/Ph.Eur.	
	TOTA	L 200.00

Pseudoephedrine Sulfate Sustained Release Layer

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and

	INGREDIENT	mg/composition
25	Pseudoephedrine Sulfate USP Hydroxypropyl Methylcellulose 2208, 1000,00c	120.0 ps
	USP/Ph.Eur.	105.0
	Microcrystalline Cellulose NF/Ph.Eur./JP	103.5
	Hydroxypropyl Methylcellulose 2910	10.5
30	Edetate Disodium	3.5
	Silicon Dioxide NF	5.0
	Magnesium Stearate NF/Ph.Eur.JP(Non-Bovin	e) 2.5

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TOTAL Tablet Weight 550.0

21) The compressed bilayer solid composition of any preceding claim wherein the total amount of desloratedine degradation products is less than about 2%.

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In ational Application No PCT/US 00/34404

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 A61K9/24 A61K A61K31/4545 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 7 A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) WPI Data, PAJ, EPO-Internal, CHEM ABS Data C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category * 1-8. EP 0 396 404 A (SCHERING CORPORATION) 10-12, 7 November 1990 (1990-11-07) 15-18,21 9,13,14, the whole document Α 19,20 page 2, line 4 - line 6 & US 5 100 675 A 31 March 1992 (1992-03-31) cited in the application 1,2,4-8, WO 99 62516 A (SCHERING CORPORATION) Y 10-12, 9 December 1999 (1999-12-09) 16-18,21 the whole document

X	Further documents are listed in the	continuation of box C.
---	-------------------------------------	------------------------

EP 0 577 957 A (J. URIACH & CIA. S.A.)

12 January 1994 (1994-01-12) page 12, line 15 - line 24

Patent family members are listed in annex.

1,3,15

 Special 	categories of	cited	documents:

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INTERNATIONAL SEARCH REPORT

In ational Application No PCT/US 00/34404

	ntion) DOCUMENTS CONSIDERED TO BE RELEVANT Challon of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
ategory *	Citation of document, with indication, where appropriate, at the	
Ρ,Υ	WO 00 02560 A (SCHERING CORPORATION) 20 January 2000 (2000-01-20) the whole document	1,3,15
1	WO 98 34614 A (SEPRACOR, INC.) 13 August 1998 (1998-08-13) page 10, line 26 -page 11, line 5	1-21
	EP 0 264 259 A (TAISHO PHARMACEUTICAL CO. LTD) 20 April 1988 (1988-04-20) the whole document	1-21
	X	

1

INTERNATIONAL SEARCH REPORT

Information on patent family members

In ational Application No
PCT/US 00/34404

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
EP 396404	A	07-11-1990	US AT AU CA DE DE DK EP ES HK JP KR WO US	4990535 A 101517 T 628986 B 5664890 A 2054752 A,C 69006628 D 69006628 T 396404 T 0471009 A 2062355 T 184896 A 6006536 B 4501425 T 9411246 B 9203278 A 9013295 A 5100675 A	05-02-1991 15-03-1994 24-09-1992 29-11-1990 04-11-1990 24-03-1994 26-05-1994 14-03-1994 19-02-1992 16-12-1994 11-10-1996 26-01-1994 12-03-1992 03-12-1994 01-07-1992 15-11-1990 31-03-1992
WO 9962516	Α	09-12-1999	US AU EP NL NL NO	6132758 A 4308599 A 1082117 A 1012191 C 1012191 A 20006079 A	17-10-2000 20-12-1999 14-03-2001 04-01-2000 03-12-1999 30-11-2000
EP 577957	A	12-01-1994	ES AT CA DE DK ES HK JP KR MX US	2042421 B 124939 T 2096318 A,C 69300255 D 69300255 T 577957 T 2076817 T 1006169 A 2730612 B 6087856 A 156518 B 9302958 A 5407941 A 5476856 A	01-08-1994 15-07-1995 23-11-1993 17-08-1995 04-01-1996 04-12-1995 01-11-1995 12-02-1999 25-03-1998 29-03-1994 16-11-1998 01-11-1993 18-04-1995 19-12-1995
WO 0002560	A	20-01-2000	AU BR EP NO	4953199 A 9910449 A 1073438 A 20005485 A	01-02-2000 02-01-2001 07-02-2001 09-03-2001
WO 9834614	A	13-08-1998	AU BR CN CZ EP HU NO PL SK ZA	6271998 A 9806157 A- 1246794 T 9901194 A 0969836 A 0001527 A 992157 A 334232 A 47299 A 9800977 A	26-08-1998 09-01-2001 08-03-2000 11-08-1999 12-01-2000 28-04-2001 04-05-1999 14-02-2000 13-03-2000 30-07-1998
EP 264259	Α	20-04-1988	JP AT	63096126 A 59294 T	27-04-1988 15-01-1991

INTERNATIONAL SEARCH REPORT Information on patent family members

in stional Application No PCT/US 00/34404

Patent document cited in search repor	t	Publication date	P	atent family member(s)	Publication date
EP 264259	A		DE US	3767114 D 4906647 A	07-02-1991 06-03-1990
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